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(54) Title: MULTIVARIATE RANDOM SEARCH METHOD WITH MULTIPLE STARTS AND EARLY STOP FOR IDENTIFICATION OF DIFFERENTIALLY EXPRESSED GENES BASED ON MICROARRAY DATA

(57) Abstract: The present invention provides multivariate methods for analyzing microarray gene expression data of high dimensional space and thereby identifying differentially expressed genes. The methods of this invention provide a random search procedure with multiple starts and early stop. Larger sets of differentially expressed genes may be identified using the methods of this invention starting from feature spaces of smaller dimensionality where accurate estimates on covariance matrix can be made.

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## MULTIVARIATE RANDOM SEARCH METHOD WITH MULTIPLE STARTS AND EARLY STOP FOR IDENTIFICATION OF DIFFERENTIALLY EXPRESSED GENES BASED ON MICROARRAY DATA

### 10 BACKGROUND OF THE INVENTION

#### FIELD OF THE INVENTION

The present invention relates in general to statistical analysis of microarray data generated from nucleotide arrays. Specifically, the present invention relates to identification of differentially expressed genes by multivariate microarray data analysis. More specifically, the present invention provides an improved multivariate random search method for identifying large sets of genes that are differentially expressed under a given biological state or at a given biological locale of interest. The method of the invention implements multiple starts and early stop in the random search of sets of differentially expressed genes.

#### DESCRIPTION OF THE RELATED ART

Gene expression analyses based on microarray data promises to open new avenues for researchers to unravel the functions and interactions of genes in various biological pathways and, ultimately, to uncover the mechanisms of life in diversified species. A significant objective in such expression analyses is to identify genes that are differentially expressed in different cells, tissues, organs of interest or at different biological states. So identified, a set of differentially expressed genes associated with a certain biological state, e.g., tumor or certain pathology, may point to the cause of such tumor or pathology, and thereby shed light on the search of potential cures.

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In practice, however, gene expression studies are hampered by many difficulties. For example, poor reproducibility in microarray readings can obscure actual differences between normal and pathological cells or create false positives and false negatives. The tension between the extremely large number of genes present (hence high dimensionality of the feature space) and the relatively small number of measurements also poses serious challenges to researchers in making accurate diagnostic inferences.

Existing methods for selecting differentially expressed genes are typically univariate, not taking into account the information on interactions among genes. As appreciated by an ordinary skilled molecular biologist, genes do not operate in isolation - activation of one gene may trigger changes in the expression levels of other genes. That is, genes may be involved in one or more pathways. Therefore, determination of differentially expressed genes calls for consideration of covariance structure of the microarray data, in addition to, for example, mean expression levels. In this regard, however, application of well-established statistical techniques for multidimensional variable selection encounters much difficulty. This is so because, in one aspect, the small number of independent samples and the presence of outliers make the estimates on selected variables unstable for large dimensions. In other words, only small sets of genes can be meaningfully considered while a relatively large number of genes are potentially differentially expressed. It is generally impossible to compare all gene subsets and find the optimal one because the number of possible gene combinations is prohibitively large. On the other hand, if a global optimum could be found, it might be overly specific to a training sample due to overfitting. Thus, it remains a significant challenge to scale methods for identifying differentially expressed genes to deal with microarray data of high dimensional space.

Therefore, there is a need to address the difficulties in applying multivariate analysis to microarray data - a need to establish rigorous methods

for identification of differentially expressed genes from high dimensional gene expression data.

#### SUMMARY OF THE INVENTION

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It is therefore an object of this invention to provide multivariate methods for analyzing microarray gene expression data of high dimensional space and thereby identifying differentially expressed genes. Particularly, it is an object of this invention to provide methods for identifying larger sets of differentially expressed genes starting from feature spaces of smaller dimensionality where accurate estimates on covariance matrix can be made. More particularly, the present invention provides a random search method with multiple starts and early stop.

In accordance with the present invention, there is provided methods for identifying a set of genes from a multiplicity of genes whose expression levels at a first and a second state, in a first and a second tissue, or in a first and a second types of cells are measured in replicates using one or more nucleotide arrays, thereby generating a first plurality of independent measurements of the expression levels for the first state, tissue, or type of cells and a second plurality of independent measurements of the expression levels for the second state, tissue, or type of cells. The method comprises: (a) identifying a quality function capable of evaluating the distinctiveness between the first plurality and the second plurality; (b) selecting a subset of genes, whose expression levels in the first and second states, tissues, or types of cells are represented in the first plurality and the second plurality, respectively; (c) calculating the values of the quality function for the subset of genes in the first state and said second state based on the first and second plurality, thereby determining the distinctiveness of the first and the second plurality; (d) substituting a gene in the subset with one outside of the subset, thereby generating a new subset, and repeating step (c), keeping the new subset if the distinctiveness increases and

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the original subset if otherwise; (e) repeating steps (c) and (d) for a first predetermined number of times, thereby identifying a locally optimal subset of genes; (f) repeating steps (b) to (e) for a second predetermined number of times, thereby identifying the second predetermined number of the locally optimal subsets; and (g) integrating the second predetermined number of the locally optimal subsets into the set of genes, wherein the set is larger than the locally optimal subsets in size.

According to the present invention, in certain embodiments, the states may be biological states, physiological states, pathological states, and prognostic states. In other embodiments, the tissues may be normal lung tissues, cancer lung tissues, normal heart tissues, pathological heart tissues, normal and abnormal colon tissues, normal and abnormal renal tissues, normal and abnormal prostate tissues, and normal and abnormal breast tissues. In yet other embodiments, the types of cells may be normal lung cells, cancer lung cells, normal heart cells, pathological heart cells, normal and abnormal colon cells, normal and abnormal renal cells, normal and abnormal prostate cells, and normal and abnormal breast cells. In still other embodiments, the types of cells may be cultured cells and cells isolated from an organism.

According to another embodiment of this invention, the integrating is performed by selecting the genes whose frequency of occurrences in the second predetermined number of the locally optimal subsets exceeds a third predetermined number. In certain embodiments, the third predetermined number is 1% or 5%. According to yet another embodiment, the first predetermined number is sufficiently small such that the global maximum is not reached. According to still another embodiment, the quality function is a parametric function or a non-parametric function. In a further embodiment, the parametric function is selected from the group consisting of the Mahalanobis distance and the Bhattacharya distance.

In various embodiments of the invention, the nucleotide arrays may be arrays having spotted thereon cDNA sequences and/or arrays having synthesized thereon oligonucleotides.

#### **BRIEF DESCRIPTION OF DRAWINGS**

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Fig. 1 depicts the steps of multivariate random search with multiple starts and early stop according to one embodiment of the invention.

Fig. 2 shows the differences of gene selection using multivariate random search with early or late stop according to various embodiments of the invention. First row are histograms of the values from the "last best iteration" in the N<sub>cycle</sub> search. Second row are histograms of the estimated Mahalanobis distances for the N<sub>cycle</sub> selected sets. Third row are histograms of the frequency of occurrences of the differentially expressed genes (1-20) in one of the selected sets.

Fig. 3 shows ROC curves for various values of N<sub>iter</sub> controlling the stopping time based on 10 simulated data sets, error bars depicting the corresponding standard errors.

Fig. 4 shows the differences of gene selection from same or different tissues using multivariate random search with early or late stop according to various embodiments of the invention. First row are histograms of the values of the "last best iteration" in the N<sub>cycle</sub> searches. Second row are histograms of the estimated Mahalanobis distances for the N<sub>cycle</sub> sub-optimal sets.

Fig. 5 shows the differences of the frequency of inclusion in the selected locally optimal set using multivariate random search according to one embodiment of the invention, applied to same or different tissue samples and with or without controls.



#### DETAIL DESCRIPTIONS OF DISCLOSURE

#### Definition

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As used herein, the term "microarray" refers to nucleotide arrays; "array," "slide," and "chip" are used interchangeably in this disclosure.

Various kinds of nucleotide arrays are made in research and manufacturing facilities worldwide, some of which are available commercially. There are, for example, two kinds of arrays depending on the ways in which the nucleic acid materials are spotted onto the array substrate: oligonucleotide arrays and cDNA arrays. One of the most widely used oligonucleotide arrays is GeneChip<sup>TM</sup> made by Affymetrix, Inc. The oligonucleotide probes that are 20- or 25-base long are synthesized in silico on the array substrate. These arrays tend to achieve high densities (e.g., more than 40,000 genes per cm²). The cDNA arrays, on the other hand, tend to have lower densities, but the cDNA probes are typically much longer than 20- or 25-mers. A representative of cDNA arrays is LifeArray made by Incyte Genomics. Pre-synthesized and amplified cDNA sequences are attached to the substrate of these kinds of arrays.

Microarray data, as used herein, encompasses any data generated using various nucleotide arrays, including but not limited to those described above. Typically, microarray data includes collections of gene expression levels measured using nucleotide arrays on biological samples of different biological states and origins. The methods of the present invention may be employed to analyze any microarray data; irrespective of the particular microarray platform from which the data are generated.

Gene expression, as used herein, refers to the transcription of DNA sequences, which encode certain proteins or regulatory functions, into RNA molecules. The expression level of a given gene refers to the amount of RNA transcribed therefrom measured on a relevant or absolute quantitative scale. The measurement can be, for example, an optic density value of a fluorescent or radioactive signal, on a blot or a microarray image. Differential expression,

as used herein, means that the expression levels of certain genes are different in different states, tissues, or type of cells, according to a predetermined standard. Such standard maybe determined based on the context of the expression experiments, the biological properties of the genes under study, and/or certain statistical significance criteria.

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The terms "vector," "probability distance," "distance," "the Mahalanobis distance," "the Euclidean distance," "feature," "feature space," "dimension," "space," "type I error," "type II error," and "ROC curve" are to be understood consistently with their typical meanings established in the relevant art, i.e. the art of mathematics, statistics, and any area related thereto. For example, a set of microarray data on p distinct genes represents a random vector  $X = X_1, \ldots, X_p$  with mutually dependent components.

#### Improved Random Search Procedure with Multiple Starts and Early Stop

Random search algorithms have been used for finding optima in complex combinatorial spaces. See, e.g., Zhigljavsky AA., Vol. 65, Mathematics and its Applications, Kluwer Academic Publishers Group, Dordrecht, 1991. The improved random search procedure according to one embodiment of this invention applies a local search procedure multiple times and then integrates the selected sets of genes to build a global optimal set of differential expressed genes. To prevent overfitting, short local searches may be performed. Local maximum regions are carefully examined and convergence to a unique global maximum is avoided. The method can be applied in conjunction with a variety of parametric and non-parametric quality functions, which are discussed in more detail in the next section. In certain embodiments, the improved random search procedure with multiple starts and early stop includes the following steps:

1. Randomly select  $N_{\text{subset}}$  genes from  $N_{\text{all}}$ , wherein  $N_{\text{subset}}$  is the number of genes in a subset,  $N_{\text{all}}$  is the total number of the genes, and  $N_{\text{subset}}$  is smaller then  $N_{\text{all}}$ .



- 2. Evaluate the quality function for the  $N_{\text{subset}}$  genes.
- 3. Generate a new evaluation point (i.e., starting point) by swapping one or more randomly selected genes between the currently selected set and the rest of the genes, thereby identifying a new  $N_{\text{subset}}$ .
- 4. Evaluate the quality function for the new  $N_{\text{subset}}$  genes; if its value has decreased, then return to the previous  $N_{\text{subset}}$ , otherwise maintain the new  $N_{\text{subset}}$ .
- 5. Repeat steps 3 and 4 until the number of iterations reaches a predetermined number let it be  $N_{\text{iter}}$  then save the resultant  $N_{\text{subset}}$  at that point.
  - 6. Repeat steps 1-5 N<sub>cycle</sub> times.

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7. Evaluate the resultant  $N_{\text{cycle}}$  groups of  $N_{\text{subset}}$  genes to identify an integrated larger set of genes.

In step 7, a post-processing step, the local optima are combined to provide a final, global solution, i.e., an integrated larger set of differentially expressed genes. Heuristically, strongly differentially expressed genes should appear in many of the local maxima. Therefore, each gene to be included in the final set of differentially may be identified based on the frequency of its occurrence in the sub-optimal (i.e., locally optimal) sets derived from each of the  $N_{\rm cycle}$  cycles, as performed in steps 1-6 above. A conservative estimate of the p-value corresponding to the observed frequency can be calculated. For example, if a gene is not differentially expressed, the probability that it will be in the selected subset by chance is expected to be equal to  $N_{\rm subset}$  /  $N_{\rm all}$ , and most likely smaller. As the number of repetitions  $N_{\rm cycle}$  is large, the final selection frequency of this gene may be approximated by a Poisson distribution with a mean  $N_{\rm cycle} \cdot N_{\rm subset}$  /  $N_{\rm all}$ . Based on this null-distribution the corresponding p-values for each gene may be calculated.

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Generally, N<sub>subset</sub> is limited by the number of available training samples (e.g., the number of microarray slides in a typical experiment) and hence, N<sub>subset</sub> may be significantly smaller than N<sub>all</sub>. Depending upon the particular quality function of choice, the nature and the extent of this limitation may vary; but, generally, both parametric and non-parametric criteria are sensitive to the scarceness of training samples in a high-dimensional feature space. In this connection, one significant advantage of the improved random search method disclosed herein is that, the detectable number of the differentially expressed genes is not limited by N<sub>subset</sub>, even though the depth of the estimated interaction structure (e.g., the covariance matrix) may be affected. In other words, a relatively large set of differentially expressed genes may be identified by integrating the subsets of genes selected from multiple local searches. In some embodiments of this invention, the final set of differentially expressed genes is significantly larger in size than the subset identified in the local search, i.e., the locally optimal subset: N<sub>subset</sub>.

The determination of  $N_{iter}$  is crucial for preventing overfitting. It cannot be too small because a small value may not permit finding truly differentially expressed genes. On the other hand, too large a number will not be efficient. When the value is too big, the same maximum may be attained in many iterations of search because of overfitting.

With regard to  $N_{cycle}$ , this is a number that substantiates the variability of this random search procedure. It may be as large as possible, only limited by the applicable CPU power (e.g.,  $N_{cycle} = 1,000,000$  may be used).

# Quality Functions Used in Conjunction with the Random Search Procedure

A variety of quality functions may be used in conjunction with the improved random search procedure in various embodiments of this invention.

A quality function measures the "distinctiveness" of the two tissues or two biological states under comparison based on a set of genes, taking into account

the correlation structure. Generally, properly specified parametric methods are more powerful than non-parametric methods due to the utilization of additional information accounted in the model, although such parametric quality functions may be sensitive to any departure from the model. With microarray data, since small sample sizes are a prevalent problem, choosing an appropriate parametric quality function may be advantageous in its power, whereas a non-parametric random search method may be more robust. One parametric measure of the differences between two multidimensional samples is the Mahalanobis distance, which is used in one embodiment of this invention. See, Mahalanobis PC., Proceedings of the National Institute of India (1936) 2 Vol. 49.

$$R_{Mah}^2 = (\mathbf{v} - \mathbf{u})' \left(\frac{\Sigma_u + \Sigma_v}{2}\right)^{-1} (\mathbf{v} - \mathbf{u}),$$

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where v and u are the sample means and  $\sum_{u}$ ,  $\sum_{v}$  are the two sample variance-covariance matrices. It is a natural extension of the *t*-statistic to a multidimensional setting. Because of the matrix inverse involved, the calculation of the Mahalanobis distance at every step of the search - for  $N_{cycle}$ .  $N_{iter}$  times - may appear to be prohibitive. However, with the improved random search procedure of this invention, changes in the vectors are only in one dimension on every step (see supra, steps 1-5); therefore, a fast update formula may be permitted. See, e.g., McLachlan GJ., Discriminant Analysis and Statistical Pattern Recognition, (1992) Wiley, NY.

In another embodiment of this invention, the Bhattacharya distance may be used, especially where differences in both the mean and the covariance structure are of interest.

$$R_{Bho}^2 = \frac{1}{8}R_{Mah}^2 + \frac{1}{2}\ln\frac{|\frac{\Sigma_u + \Sigma_v}{2}|}{\sqrt{|\Sigma_v||\Sigma_v|}}.$$

Similarly, other parametric or non-parametric dissimilarity measures may be used in various alternative embodiments in conjunction with the

improved random search procedure disclosed herein. Such different choices of quality functions each may be designed to deal with microarray data with different characteristics.

Further, when using various quality functions, various background reduction, normalization, and other adjustment procedures may be applied to the microarray data. For example, rank-based adjustment and the typical mean-log adjustment (dividing by mean and take logarithm) may be used. In one embodiment, the following adjustment is implemented: the data points on each slide or array were replaced by their normal scores using the formula

 $X_{ij}^{n} = \Phi^{-1}(\operatorname{rank}_{i}X_{ij}/N_{all}),$ 

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where  $\Phi^{-1}(\alpha)$  is the  $100\alpha^{th}$  percentile of the standard normal distribution and  $\operatorname{rank}_j \mathbb{X}_{ij}$  is the rank of  $\mathbb{X}_{ij}$  among all of the observations on the  $j^{th}$  slide. See, Tsodikov A. et al., (2002) Bioinformatics 18: 251-260.

#### Computer Simulation of the Multivariate Search Method

A simulation study was performed to evaluate the improved random search method. Totally 1000 genes were divided into subsets of equal size 20. One of the subsets was selected to be deemed as differentially expressed with the gene-specific ratio d randomly generated for each of the genes from a lognormal distribution with mean 1 and variance 0.5. The correlation structure was kept the same in the two hypothetic tissues. In the selected subset some of the genes exhibited large over- or under-expression, while others with  $d \approx 1$  changed their expression level only slightly. The simulation was performed on 20 slides or arrays with one of the tissues on the green channel and the other on the red channel. The relevant parameters for the random search were set:  $N_{cycle} = 10,000$ ,  $N_{subset} = 5$ ; and, the Mahalanobis distance was used as the quality function.

Referring to Fig. 1, the results of the random search procedure are compared between  $N_{iter} = 1000$  (in the left panels) and  $N_{iter} = 100,000$  (in the right panels). The two graphs on the top show the histograms of the values of the "last good iteration" - the number of iterations after which no new successful steps were encountered (i.e., when no new subset was found any more at step 4 of the aforementioned procedure and thus the final set was determined). The two histograms demonstrate that 1000 iterations were a little less than sufficient to reach the global maximum, whereas 10,000 iterations were more than enough for the random search to converge.

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The middle graphs illustrate the same phenomenon in another way. In the case of early stopping, i.e., when  $N_{iter} = 1000$ , the distribution of the Mahalanobis distances corresponding to the  $N_{cycle}$  sub-optimal sets is unimodal with high variability. Thus, the procedure has explored many different local maxima with a variety of corresponding values of the quality function. On the other hand, when the number of iterations increase, e.g., when  $N_{iter} = 100,000$ , the distribution of the Mahalanobis distances achieved in the sub optimal sets became very discrete. In about half of the cases the search reached the global maximum on a unique combination of genes. Therefore, in this situation, although the global maximum was found, many local maxima and the corresponding differentially expressed genes from the various subsets were missed. When early stop is carried out at the 1000-th iteration, none of 10,000 cycles found the global maximum, but a variety of genes were selected.

At the bottom panel of Fig. 1, the frequencies of selection for the 20 genes in the differentially expressed gene set are plotted. The x-axis represents the number of the genes: from gene No. 1 to No. 20. With  $N_{iter} = 1,000$ , i.e., when the early stop was implemented, 17 from 20 genes pass the selection criteria (predetermined to be a frequency of occurrence higher than 0.5%). With  $N_{iter} = 100,000$ , i.e., when the early stop was not implemented, only 10 genes met the 0.5% frequency standard when the global maximum was attained.

Referring to Fig. 2, the ROC curves corresponding to values of Niter ranging from 100 to 10,000 based on 10 independently simulated data sets were plotted. Other parameters were held constant, that is,  $N_{subset} = 5$ ,  $N_{cycle} =$ 10,000. For each search, a list of genes with associated frequencies of occurrence in the selected subsets were complied and a final set of differentially expressed genes was identified by applying cutoff values ranging from 0.1% to 10% in frequency. Based on the null hypotheses of no differential expression, for each of these sets, the ratio of type I error (i.e., the false positive) was defined as the proportion of non-differentially expressed genes that was selected into the final set. And the ratio of the type II error (i.e., the false negative) was defined as the proportion of the genes in the differentially expressed subsets that were not included in the final set. The resulting ROC curves are shown in Fig. 2. Also, as a reference, the point representing the type I error and the power of the marginal t-test with 5% significance level is also plotted (referring to the star in Fig. 2). Comparing the ROC curves in Fig. 2, a skilled artisan can note that the value of Niter significantly affects the performance of the random search procedure: long searches are inferior to searches with early stop. There ought to be, however, a limit on how early the search should stop, because very short searches are not likely to reach any local maxima. According to Fig. 2, the best performance was achieved when  $N_{iter} = 500$ .

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The invention is further described by the following examples, which are illustrative of the invention but do not limit the invention in any manner.

# Example 1: a Detailed Illustration of Random Search with Multiple-Starts and Early Stop

Referring to Fig. 3, suppose there are p genes and n and m independent samples in the two classes respectively, this procedure finds a group of genes differentially expressed in these classes using information on the k-variate dependence structure.



1. Repeat the following  $N_{iter}$  times.  $N_{iter}$  is not too large; early stop – stop before convergence – is implemented.

- a. Randomly select k genes (genes 2 to gene k in Fig. 3) that will serve as the seed of the random search.
- b. Calculate the distance between the two classes based on the k initially selected genes.
- c. Randomly select a gene (e.g., gene 2 in Fig. 3) from the current gene set (gene 2 to gene k in Fig. 3), remove it from the set and replace it with a gene randomly selected from outside of the set (e.g., any of gene k+1 to gene p in Fig. 3, let it be gene x).
- d. Calculate the distance between the two classes based on the new gene set (gene 3 to gene k, plus gene x). If this distance is larger than the previously calculated one, then keep the change, otherwise revert to the previous set.
- e. Retain the selected sub optimal set of genes, i.e., the set that has the largest distances between the two classes.
  - 2. Repeat step 1  $N_{\text{cycle}}$  times, obtain  $N_{\text{cycle}}$  sets of genes of size k.
  - 3. For each gene, calculate the frequency of its occurrence as a member of a sub-optimal set.
- 20 4. The final set of genes is defined as the genes that have a frequency of occurrence exceeding a preset limit.

# Example 2: a Source Code Segment Implementing Random Search with Multiple Starts and Early Stop - Step 1 and 2 of Example 1

25 Program gene1

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c parameter (nall=1000, ncl=10, niter=500, m=20,l=2,nt=2) parameter (ishift=3000,NCYCLE=1000) parameter (genadd=5.,disp=1.,debug=2.)

```
parameter (exprnax=20.,strang=1.e-15)
        parameter (kcl=5,iap=1,nex=10)
        parameter (pat=1.5,dpat=0.,frailty=0.2,ncls=20,purity=0.85)
        CHARACTER*50 jmode,qualit, ranf,ku,stat,start,normal,mixup
        CHARACTER*50 sound,ill
        DIMENSION AP(L*IAP), DEL(M*I)
        DIMENSION DEN((KCL+2)*L),PST(L),DFM(L*(KCL+2)*L*iap)
        DIMENSION F(KCL+2), DS(M*L*L*(KCL+2))
        DIMENSION DI(ncl), DETER(L), rank1(m), rank2(m)
 10
        dimension err(kcl+2),g((kcl+2)*1),ent(l)
       Dimension inum(ncl),b(nall*m*l),a(nall*m*l),cl(ncl*m*l),u(m*l)
       dimension e(ncl*ncl),ito(l),ind3(niter)
        dimension e1(ncl*ncl),e2(ncl*ncl),e3(ncl*ncl),z(nex*nex)
       dimension imbest(ncl),x(m*1),v(nall),m22(m*1),ind2(nall)
       dimension r(ncl*ncl*l),r2(ncl*l),r3(ncl*ncl*l)
       dimension mv(kcl),ff(kcl),dd(kcl),rr(kcl)
       dimension stud(nall), tkolm(nall), tmann(nall)
       dimension iex(nex)
       character*10 ndata, ntime
       data iex/1,2,3,4,5,6,7,8,9,10/
       data f/0.5,0.6,0.7,0.8,0.9,1.0,1.1/
 25
       data ap/0.5,0.5/
       data qualit /'mahalo'/
       data imode /'one-leave-out-'/
       data mixup /'no'/
30
       data ranf / ffile'/
       data normal/'gauss'/
       data stat /'param'/
       data start / bestcor'/
35
       sound='red1.txt'
       ill='red2.txt'
       OPEN (unit=NT,FILE='b', FORM='FORMATTED',STATUS='unknown')
       open(unit=11,file=sound,form='formatted',
40
       * status='old')
       open(unit=22,file=ill,form='formatted',
       * status='old')
       open(unit=68, file='inbest.dat', form='formatted',
       * status='unknown')
45
       open(unit=69,file='best.dat',form='formatted',
       * status='unknown')
       write(nt,'(//30x,"GENE CLUSTER MASTER"/)')
       write(nt,'("Number of slides M = ", i3)')m
50
       write (nt, "Number of genes NALL =", i5,
       * ",gemuin cluster size=",i4,",to be searched:",i4)")
       * nall,ncis,ncl
       write (nt,'("DATA normalization to(by) = ",A10)')normal
       write (nt,'("Type of Statistics Used = ",A10)')stat
55
       if(ranf.ne.'ffile')then
       write(nt,'("Overexpression of Poisoned Genes ",f5.1,
       * " Variance ",f5.1)")
       * genadd,disp
       write(nt,'("Random Numbers Generator ",a10," ,Shift",i5)')
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       * ranf,ishift
```

```
end if
         write(nt,'(//30x,"SIMULATION PARAMETERS"/)')
         write(nt,'("Sound data from: "a30)')sound
         write(nt,'("Patology data from: "a30)')ill
         write(nt,'("SIMULATED PATOLOGY LEVEL: ",f3.1,"+/-",f3.1)")
         write(nt,'("Level of mutual Frailty for Cluster: ", f5.2)")
         * frailty
         write(nt,'("Mixture: ",f5.2,"LogNorm+",f5.2"Uniform")')
 10
         * purity,1.-purity
        write(nt,'(//30x,"SEARCH PARAMETERS"/))
         write(nt,'("MIXUP the GENES? ", a10)")mixup
         write(nt,'("SEARCH MODE ", a10)')qualit
 15
        write(nt,'("Number of Random Search Trials:"i7)")
        * niter
        if (nex.ne.0) then
        write(nt,'("ATTENSION!, Genes Excluded:"/10(10i6/))')
        * (iex(i),i=1,nex)
 20
        end if
        if(qualit.eq.'parz'.or.qualit.eq.'km')then
        write(nt,'("MODE OF BAYES QUALITY ", a10)')ku write (nt,'("Number and values of kernels",i5/
        * 15 f5.1)')kcl,(f(i),i=1,kcl)
 25
        end if
        do i=1,ishift
        aa=mdm(-1.)
        end do
30
        if(ranf.eq.'uni')then.
        do i=1,nali*m*2
        b(i)=1.+mdm(-1.)*disp
        end do
35
        else if (ranf.eq.'normco')then
        do i=1,nall*m-1,2
       call normco(b(i),b(i+1),5.,3.,disp,disp,0.9)
       end do
40
       do i=nall*m+1,nall*m*2-1,2
       call normco(b(i),b(i+1),5.,3.,disp,disp,-0.9)
       end do
       else if(ranf.eq.'ffile') then
       call rfromf(b, nall,m,l)
45
       C
       else
       write(nt,'("no such data mode",a10)")ranf
       stop 67
       end if
50
       if(ranf.ne.'ffile') then
       do j=m,2*m-1
       do i=nall*j+1,nall*j+ncl
55
       b(i)=b(i)+genadd
       end do
       end do
       if (nall.le.10) then
       write (nt,'(10f7.2/)')b
      end if
```

```
end if
         if(mixup.eq.'yes') then
         do i=1,nall
         ind2(i)=i
         end do
         do i =1, ishift
         iin=mdm(-1.)*nall+1
         iout=mdm(-1.)*nall+1
  10
        numold=ind2(iout)
         ind2(iout)=ind2(iin)
        ind2 (iin)=numold
        if(iin.gt.nall.or.iout.gt.nall.or.iin.lt.1.or.iout.lt.1) then
        write(*,'("BIGGGGG!!!!", 3i18)')i,iin,iout
 15
        end if
        call exchange(b,nall,m,l,iin,iout,x,u)
        end do
        if (debug.ge.5) then
        write (nt,'("Mixed Cluster"/1000(10i8/))')
 20
        * (ind2(i),i=1,nall)
        end if
        end if
        if (normal.ne.'no') then
 25
        call normalization(b,ind2,nall,m,l,stud,tkolm,normal)
        call tests(b,m22,ind2,nall,m,l,x,u,stud,tkolm,tmann,nt,ncl)
 30
       ito(1)=m
       ito(2)=m
       mb=ito(1)+ito(2)
       istg=0
35
       sd=0.
       stiter=1.e20
       do i=1,m*1
       u(i)=1./m
40
       end do
       if(start.eq.'bestcor'.and.nex.ge.2) then
       do i=1,nex
       inum(i)=iex(i)
45
       end do
       call assign(b,inum,cl,nex,nall,m,l)
       c write (*,'("u(i) "/10(10f8.5/))")
       c * (u(i), i=1,m*l)
50
       call misr1(cl,r2,r,u,nex,ito,mb,l)
       call cover(r,r3,z,nex,m,l)
       write (nt,'(/25x,"CORRELATION MATRIX"/10(12i6/))')
       * (iex(i), i=1,nex)
       write (nt,'(/10(10f6.2/))')
55
       * (r3(i), i=1,2*nex*nex)
       write (nt,'(/25x,"FISHER MATRIX"/10(10i6/))')
      * (iex(i), i=1,nex)
      write (nt, '(/10(10f6.2/))')
60
      * (z(i), i=1,nex*nex)
```

```
write (nt,'(/"Genes means "/5(10f6.2/))")
        * (r2(i), i=1,2*nex)
        call bhafas(r,r2,e,e1,e2,e3,rb,rm,rc,nex,qualit,debug)
  5
        write (nt,'(/"Mahalonobis Distance: ",f12.2)')rm
        stop 777
        end if
       DO ICY=1,NCYCLE
 10
       ii=0
        if(start.eq.'random') then
       iin-mdm(-1.)*nall+1
       imm(1)=iin
 15
       do i=2,ncl
       88 continue
       inew=rndm(-1.)*nall+1
       do j=1,i-1
 20
       if(inew.eq.imm(i-j))then
       go to 88
       else
       imm(i)=inew
       end if
 25
       end do
       end do
       else if(start.eq.'last') then
       DO I=1,NCL
       ii=ii+1
30
       inum(ii)=i+NALL-NCL
       end do
       else if(start.eq.'first') then
       do i=nall, nall-ncl,-1
       ii=ii+1
       inum(ii)=i+NALL-NCL
       end do
       else if(start.eq.'frombest') then
       read(68,'i7,e12.4,(10(10i6/))')ll,qq,(inum(i),i=1,ncl)
       else
40
       stop 9999
       end if
       write (nt,'(" Initially Selected genes "/
       * 5(10i5/))')inum
45
       DO iter=1,niter
       if (iter.ne.1)then
       call change(inum,nall,ncl,iin,iout,numold,ind3,iter,niter,
       * iex,nex)
50
       else
       iin=1
      iout=1
       mmold=99
       if (iter.gt.1.and.iter.le.5.and.debug.ge.3.) then
       write (nt, '("Iteration", i4," Exchanged genes ", 3i5/
      * "MASK Array"/
      * 5(10i5/))')iter,IIN,iout,numold,(inum(i),i=1,ncl)
60
```

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```
call assign(b,inum,cl,ncl,nall,m,l)
         if(stat.eq.'param')then
        call misr1(cl,r2,r,u,ncl,ito,mb,l)
        if (debug.ge.3) then
write (nt,'("Genes cov "/5(5f12.5/))")r
call cover(r,r3,z,nc1,m,1)
 10
        write (nt, '("Genes cor "/5(5f12.5/))')r3
        write (nt,'("Genes means ",5(5f12.2/))')r2
        end if
        else if (stat.eq.'nonparam')then
        call SPIR1(cl,r2,R,X,V,NCL,M,L,m22,ind2,rank1,rank2)
 15
        if (debug.ge.5) then
        write (nt,'("Genes spirmen "/5(10f12.5/))')r
        write (nt,'("Genes medians ",5(10f12.2/))")r2
write (nt,'("Genes interQU ",5(10f12.2/))")
        * (v(i),i=1,ncl*l)
 20
        c stop 777
        end if
        end if
        c BHATTACHARYA DISTANCE
25
        if(qualit.eq. bhata') then
        else if (qualit.eq.'mahalo') then
        ss=rm
        else if (qualit.eq.'corcor') then
30
       ss=rc
       else
       write(nt,'("no such quality function",a10)') qualit
       stop 67
       end if
       end if
       IF(SS.GT.SD) THEN
       SD=SS
40
       ISTG=ISTG+1
       c REMEMBERING OF BEST VALUES
       c CALL UCOPY(inum,imbest,ncl)
       do iu=1,ncl
       imbest(iu)=inum(iu)
       end do
       ibest=iter
50
       CALL DATIMH(NDATA,NTIME)
      WRITE(*,'("SUCCESS at: ",A12,2X,A12,
* "ITERATION", i7," QUALITY",e14.6)')
* NDATA,NTIME,ITER,SD
      write(*,'(10(10i5/))') (inum(i),i=1,ncl)
      if(debug.ge.2)then
      WRITE(nt,'("GOOD!!!,Iteration and Q",i7,e14.6)')ITER,SD
      write(nt,'(10(10i5/))') (inum(i),i=1,ncl)
      ELSE IF(SS.LE.SD) THEN
```

```
inum(iout)=numold
         if(debug.ge.3.and.iter.le.10)then
         CALL DATIMH(NDATA, NTIME)
         WRITE(nt,'("BAAD!!!,Qnew and Qbest",i7,2e14.6)')ITER,SS,SD
         end if
         END IF
         if(SD.GT.STITER) then
         write (NT, ("REQUIRED DISTANCE ACHIEVED!", 2e15.3)")
  10
         * SD, STITER
        go to 18
        end if
        END DO
        18 continue
        write(nt,'(25x," CYCLE N " i6/)')ICY
        write(nt,'("Distance used: ",A6," Quality=",e15.3)")
        write(nt,'("Number of successful steps: ",i5)")istg
write(nt,'("Best Cluster Obtained After: ",
  20
        * i9/20(10i7/))')
        * niter,imbest
        c rewind 68
        write(69,'(i7,f12.4,20i6)') ibest,sd,imbest
        c write(69,'(i7,f12.4)') ibest,sd
        END DO
        stop
        end
 30
       subroutine tests(b,m22,ind2,nall,m,l,x,u,stud,tkolm,tmann,nt,ncl)
       dimension b(nall,m,l),stud(nall), tkolm(nall),tmann(nall)
       dimension x(m*1),u(m*1),m22(m*1),ind2(nall)
       i34=m*0.75
       i14=m*0.25
       i5 = m*0.5+1
       do i=1,nall
       do j=1,m
       x(j)=b(i,j,1)
       u(j)=b(i,j,2)
40
       end do
       call sortzv(x,m22,m,1,1,0,0)
       xd=(x(m22(i34))-x(m22(i14)))/1.35
       xm=x(m22(i5))
45
       call sortzv(u,m22,1,m,1,0,0)
      ud=(u(m22(i34))-u(m22(i14)))/1.35
      um=u(m22(i5))
      stud(i)=abs(xm-um)/sqrt(xd*xd+ud*ud)
      call sortzv(stud,ind2,nall,1,1,0,0)
50
      write (nt,'("N.Student Cluster"
      */1000(10i8/))')
      *(ind2(i),i=1,ncl)
      call errors(ind2,nt,ncl,nall)
55
      do i=1,nall
      xm=0.
      xm2=0.
      um=0.
      um2=0.
```

```
do j=1,m
        x(j)=b(i,j,1)
        u(j)=b(i,j,2)
        end do
        do j=1,m
        xm=xm+x(j)
        xm2=xm2+x(j)*x(j)
        um=um+u(j)
        um2=um2+u(j)*u(j)
 10
       end do
       xm=xm/m
       um=um/m
       xd=xm2/m-xm*xm
       ud=um2/m-um*um
       stud(i)=abs(xm-um)/sqrt(xd+ud)
       end do
       call sortzv(stud,ind2,nall,1,1,0,0)
       write (nt, ("Param Student Cluster"/1000(10i8/))")
       * (ind2(i),i=1,ncl)
       call errors(ind2,nt,ncl,nall)
 20
       do i=1,nall
       do j=1,m
       x(j)=b(i,j,1)
       x(j+m)=b(i,j,2)
 25
       end do
       CALL UTEST(x,u,m,m,tmann(i),ZU,IERR)
       call sortzv(tmann,ind2,nall,1,0,0,0)
       write (nt,'("Mann-Whitney Cluster"/1000(10i8/))')
       * (ind2(i),i=1,ncl)
       call errors(ind2,nt,ncl,nall)
       do i=1,nall
       do j=1,m
       x(j)=b(i,j,1)
      u(j)=b(i,j,2)
35
       end do
      CALL kolm2(x,u,m,m,tkolm(i),Prob)
      call sortzv(tkolm,ind2,nall,1,1,0,0)
      write (nt, '("Kolmogorov Cluster"/1000(10i8/))')
       * (ind2(i),i=1,ncl)
      call errors(ind2,nt,ncl,nall)
      return
      end
45
```

## Example 3: a Source Code Segment Implementing Integration of The Results from Local Searches to Build a Larger Set of Genes - Steps 3 and 4 Of Example 1

```
Program genecount
c

55 parameter (nall=1000, nclust=5, ntrial=10000,ncut=10,nr=22,nt=2)
parameter (nctrue=20,ipat=1,ntupw=1,ntidw=17,memw=100000)
parameter (debug=2.)
c
dimension a(nclust*ntrial),c(nall),cut(ncut),genprop(nclust)

60 dimension sel(nall)
```

```
dimension tontuple(nclust+3), ind(nall,nall), ind1(nall)
        character*30 selgen
        character*8 mode
        data cut/0.000005,0.00001,0.00005,0.001,0.002,0.003,0.01,0.03,
        * 0.05,0.08/
        data cutpair/0.1/
        data cpair/0.003/
        data selgen /'best.dat'/
        data mode/'sim'/
 10
        data niter /500000/
        CHARACTER*1 opmo
        CHARACTER*50 hbname
        CHARACTER*8 tek(nclust+3)
 15
        DATA opmo/'X'/,LRECLR/1024/,LRECLW/1024/
        OPEN (UNIT=NT,FILE='b.count', FORM='FORMATTED',STATUS='UNKNOWN')
        open(unit=nr,file=selgen,form='formatted',status='old')
 20
       hbname='genome.hbook'
       tek(1)='lastb'
tek(2)='quality'
tek(3)='N_of_gen'
        tek(4)='gene1'
       tek(5)='gene2'
tek(6)='gene3'
tek(7)='gene4'
tek(8)='gene5'
 25
30
       c tek(i)='gene'//ichar(i-2)
       c end do
       if(ntupw.gt.0) then
       call HROPEN(ntidw, 'ani98', hbname, 'N', lreclw, ISTAT)
35
       call HBOOKN(ntidw, 'GENE SELECTION', nclust+3,
       * '//ani98',memw,tek)
       write(nt,'(/10x,"GENE SORTER FOR ", A10,
       * "EARLY STOP AT",18)')mode, NITER
       qmean=0.
40
       nmean=0
       ntrj=0
       ncount=1
       do i=1,nclust
       genprop(i)=0.
      end do
       do j=1,ntrial
      read(22,*,err=100,end=99)nlast,quality,
       * (a(i),i=ncount,ncount+nclust-1)
      tontuple(1)=nlast
50
      tontuple(2)=quality
      jj=-1
      kk=0
      do i=4,nclust+3
      jj=jj+1
      tontuple(i)=a(ncount+jj)
      if(tontuple(i).le.nctrue) then
      genprop(i-3)=genprop(i-3)+1.
      kk=kk+1
      end if
60
      end do
```

```
tontuple(3)=kk
         call HFN(ntidw,tontuple)
         ncount=ncount+nclust
         ntrj=ntrj+1
         nmean=nmean+nlast
         qmean=qmean+quality
         end do
         go to 99
         100 continue
 10
         write(*,'("ERROR IN INPUT STREAM ON LINE: ",i7)')j
        stop
        99 continue
        write (nt, '(i7," Random Starts, Rm and Last ",f12.4,i7)')
 15
         * ntrj,qmean/ntrj,nmean/ntrj
        if (mode.eq.'sim') then
        write (nt,'(/"% Of True",10(5f12.4/))')
* (genprop(i)/ntrj, i=1,nclust)
 20
        end if
        call vzero(c,nall)
        do i=1,nclust*ntrj
        do k=1,nall
25
        realk=real(k)
        if(a(i).eq.realk) then
        sel(k)=sel(k)+1
        c(k)=c(k)+1./ntrj
end if
30
        end do
        end do
        do j=1,ncut
        do i=1,nall
        indl(i)=0
35
        end do
        ncount=0
        do i=1,nall
       if (c(i).gt.cut(j))then
       ind1(i)=1
       if (debug.ge.3) then
40
       write(nt,'("GENE ",15,5x,
* "Appearance % ",F12.5)')I,C(I)
       end if
       ncount=ncount+1
45
       END IF
       end do
       err1=0.
       eπ2=0.
50
       do_i=1,nall
       if(ind1(i).eq.1.and.i.le.nctrue)then
       errl=errl+1.
       else if(ind1(i).eq.1.and.i.gt.nctrue)then
       еп2=еп2+1.
55
       end if
       end do
      write(nt,'(/"N of genes selected with CUT ",F9.5,i8)")
       * cut(j),ncount
       if (mode.eq.'sim') then
60
      write(nt,'("1 error: ",F9.5,", 2 error",F9.5)')
```

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```
* 1.-err1/nctrue,err2/nall
        write(nt,'("Eta = 1.- 1error/sqrt(2error): ",F12.5)')
        * err1/nctrue/sqrt(err2/nall)
        end if
        end do
        if (debug.ge.4.) then
        ncount=0
        do i=1,nall
        do j=1,nall
 10
        ind(i,j)=0
        end do
        end do
        do ni=1,ntrj
        do j=1,nclust-1
 15
       k1=ifix(a(ncount+j))
       do i=j+1,nclust
       k2=ifix(a(ncount+i))
       ind(k1,k2)=ind(k1,k2)+1
       end do
20
       end do
       ncount-ncount+nclust
       end do
       do i=1,nall
25
       do j=i+1,nall
       prop=real(ind(i,j))/sel(i)
       if(prop.ge.cutpair.and.c(i).ge.cpair)then
       write(nt, '("Freq. for genes:",216,3f12.5)')
30
       c * "Single frequencies:",9x,2f12.5)")
       * i,j,prop,c(i),c(j)
       end do
       end do
35
      end if
      if(ntupw.gt.0) then
      call HROUT(0,ICYCLE,'')
      call HREND('ani98')
40
      end if
      STOP
      END
```

50

# 45 Example 4: Microarray Expression Analysis Using Cells from Two Colon Cancer Cell Lines

HT29 cells represent advanced, highly aggressive colon tumors. They contain mutations in both the APC gene and p53 gene, two tumor suppressor genes that frequently mutate during colon tumorigenesis. HCT116 cells manifest less aggressive colon tumors and harbor functional p53 and APC. They are defective in DNA repair. The experiment was performed with three RNA samples (1 µg RNA each). Cy-3-dCTP (green) was used to label

HCT116 cells while Cy-5-dCTP (red) was used for HT29 cells. Each comparison set was hybridized against two microarray slides (facing each other) containing 4608 minimally redundant cDNAs spotted in duplicate. As control, six *Drosophila* genes were added to the Cy-5 samples. Thus, in a red vs. green comparison they are differentially expressed by design. This experiment resulted in a total of twelve measurements on each channel for each gene on the microarrays. Although a nested dependence structure existed in the samples, the analysis assumes them as independent replicates. Additionally, ten HCT116 samples hybridized with Cy-5 (red) from a separate experiment were included in the analysis.

Two comparisons were performed: (i) HCT116 vs. HT29 and (ii) HCT116 (green) vs. HCT116 (red); the first is inter cell lines whereas the second is intra cell lines. The relevant parameters for the random search were set:  $N_{cycle} = 10,000$ ,  $N_{subset} = 5$ ; and, the Mahalanobis distance was used as the quality function.

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Referring to Fig. 4, the left panel corresponds to the comparison of the different cell lines (as the case (i) above) whereas the right panel to the comparison of the same cell line on different channels (as the case (ii) above). The histograms of the last best iteration (the top two graphs) are very similar in both cases; neither has reached the global maximum. That is, in both cases, the procedure kept exploring the local maxima due to the early stopping. However, turning to the bottom two graphs, the distribution of the estimated Mahalanobis distances at these local maxima in each case is very different from each other: When different cell lines were compared, i.e., in the case (i) above, the Mahalanobis distances based on the locally optimal subsets tended to be much larger than those in the case (ii) above when the same cell lines were compared. Therefore, the separation of the two tissues was considerably better in case (i) than in case (ii), as one would expect.

Referring to Fig. 5, the first 115 genes ordered according to the decreasing frequency of occurrence in the selected subsets are plotted. The white columns represent genes from same cell line samples without control whereas the black columns represent genes from different cell line samples. In addition, the gray columns represent genes from same cell lines samples with control. As shown, the right tails of the histograms are very close to each other. Some of the genes in the HCT116/HT29 comparison (the black columns) are selected more often – i.e., have higher frequency – than expected under the null hypothesis of no difference between the two tissues (the white columns). Interestingly, in the case with same cell line without control (the white columns), only two genes had a frequency that was higher than 3%; and, when the control genes were included (the gray columns), this number increased to six and four out of the top five genes (Nos. 1, 2, 3, and 5 on the x axis) were actually *Drosophila* control genes.

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A frequency level of 1% was selected as the cutoff for identifying differentially expressed genes. Total 59 genes were selected and thus 59 cDNA spots were identified on the slides. A comparison was carried out between the 59 cDNA spots and the top 59 genes selected by t-statistic. Almost half of those genes (25 to be exact) were identified by both methods. However, a characteristic advantage of the multivariate random search procedure was its ability to identify correlated genes. Some of the genes had several corresponding spots on the slides, and therefore their expression levels at various spots were known to be correlated. Among the 59 genes identified by the multivariate random search method, 13 had two, and two had three spots inter-related to each other. By comparison, among the genes identified by the marginal t-statistic, 17 genes had two or more replicates on the slides, and only one of them had all of its replicates selected in the resulting list of genes. Therefore, the improved random search procedure of this invention is powerful in identifying less pronounced differentially expressed genes when they are correlated with more strongly differentially expressed genes.

It is to be understood that the description, specific examples and data, while indicating exemplary embodiments, are given by way of illustration and are not intended to limit the present invention. Various changes and modifications within the present invention will become apparent to the skilled artisan from the discussion, disclosure and data contained herein, and thus are considered part of the invention.

#### **CLAIMS**

20

- 1. A method for identifying a set of genes from a multiplicity of genes whose expression levels at a first state and a second state are measured in replicates using one or more nucleotide arrays, thereby generating a first plurality of independent measurements of the expression levels for said first state and a second plurality of independent measurements of the expression levels for said second state, which method comprises:
- (a) identifying a quality function capable of evaluating the distinctiveness between the first plurality and the second plurality;
  - (b) selecting a subset of genes, whose expression levels in said first state and second state are represented in said first plurality and said second plurality, respectively;
- (c) calculating the values of the quality function for said subset of genes in said first state and said second state based on the first and second plurality, thereby determining the distinctiveness of the first and the second plurality;
  - (d) substituting a gene in said subset with one outside of said subset, thereby generating a new subset, and repeating step (c), keeping the new subset if the distinctiveness increases and the original subset if otherwise;
  - (e) repeating steps (c) and (d) for a first predetermined number of times, thereby identifying a locally optimal subset of genes;
  - (f) repeating steps (b) to (e) for a second predetermined number of times, thereby identifying said second predetermined number of the locally optimal subsets; and
  - (g) integrating said second predetermined number of the locally optimal subsets into said set of genes, wherein said set is larger than said locally optimal subsets in size.

2. The method of claim 1, wherein said states are selected from the group consisting of biological states, physiological states, pathological states, and prognostic states.

- 3. A method for identifying a set of genes from a multiplicity of genes whose expression levels at a first tissue and a second tissue are measured in replicates using one or more nucleotide arrays, thereby generating a first plurality of independent measurements of the expression levels for said first tissue and a second plurality of independent measurements of the expression levels for said second tissue, which method comprises:
- (a) identifying a quality function capable of evaluating the distinctiveness between the first plurality and the second plurality;

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- (b) selecting a subset of genes, whose expression levels in said first tissue and second tissue are represented in said first plurality and said second plurality, respectively;
- (c) calculating the values of the quality function for said subset of genes in said first tissue and second tissue based on the first and second plurality, thereby determining the distinctiveness of the first and the second plurality;
- (d) substituting a gene in said subset with one outside of said subset, thereby generating a new subset, and repeating step (c), keeping the new subset if the distinctiveness increases and the original subset if otherwise;
- (e) repeating steps (c) and (d) for a first predetermined number of times, thereby identifying a locally optimal subset of genes;
- (f) repeating steps (b) to (e) for a second predetermined number of times, thereby identifying said second predetermined number of the locally optimal subsets; and

(g) integrating said second predetermined number of the locally optimal subsets into said set of genes, wherein said set is larger than said locally optimal subsets in size.

- 4. The method of claim 3, wherein said tissues are selected from the group consisting of normal lung tissues, cancer lung tissues, normal heart tissues, pathological heart tissues, normal and abnormal colon tissues, normal and abnormal renal tissues, normal and abnormal prostate tissues, and normal and abnormal breast tissues.
- 5. A method for identifying a set of genes from a multiplicity of genes whose expression levels in a first type of cells and a second type of cells are measured in replicates using one or more nucleotide arrays, thereby generating a first plurality of independent measurements of the expression levels for said first type of cells and a second plurality of independent measurements of the expression levels for said second type of cells, which method comprises:
  - (a) identifying a quality function capable of evaluating the distinctiveness between the first plurality and the second plurality;

- (b) selecting a subset of genes, whose expression levels in said first type of cells and said second type of cells are represented in said first plurality and said second plurality, respectively;
- (c) calculating the values of the quality function for said subset of genes in said first type of cells and said second type of cells based on the first and second plurality, thereby determining the distinctiveness of the first and the second plurality;
- 25 (d) substituting a gene in said subset with one outside of said subset, thereby generating a new subset, and repeating step (c), keeping the new subset if the distinctiveness increases and the original subset if otherwise;

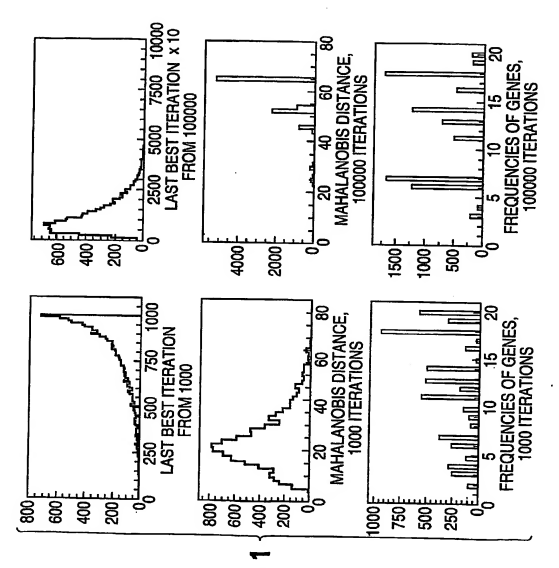
(e) repeating steps (c) and (d) for a first predetermined number of times, thereby identifying a locally optimal subset of genes;

(f) repeating steps (b) to (e) for a second predetermined number of times, thereby identifying said second predetermined number of the locally optimal subsets; and

- (g) integrating said second predetermined number of the locally optimal subsets into said set of genes, wherein said set is larger than said locally optimal subsets in size.
- 6. The method of claim 5, wherein said types of cells are selected from the group consisting of normal lung cells, cancer lung cells, normal heart cells, pathological heart cells, normal and abnormal colon cells, normal and abnormal renal cells, normal and abnormal prostate cells, and normal and abnormal breast cells.
- 7. The method of claim 5, wherein said type of cells are selected from the group consisting of cultured cells and cells isolated from an organism.
  - 8. The method of claim 1, 3, or 5, wherein said integrating is performed by selecting the genes whose frequency of occurrences in said second predetermined number of the final subsets exceeds a third predetermined number.
- 20 9. The method of claim 8, wherein said third predetermined number is 1% or 5%.
  - 10. The method of claim 1, 3, or 5, wherein said first predetermined number is sufficiently small such that the global maximum is not reached.
- 11. The method of claim 1, 3, or 5, wherein said quality function is a parametric function or a non-parametric function.

12. The method of claim 11, wherein said parametric function is selected from the group consisting of the Mahalanobis distance and the Bhattacharya distance.

13. The method of claim 1, 3, or 5, wherein the nucleotide arrays are selected from the group consisting of arrays having spotted thereon cDNA sequences and arrays having synthesized thereon oligonucleotides.



F. 6.

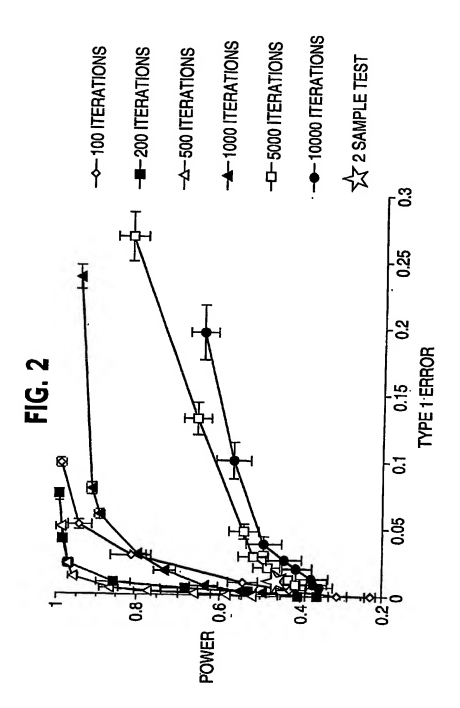


FIG. 3

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CLASS 2								
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	<b>S</b> 5							
	<b>S1</b>							
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CLASS 1

S1 S2 S3

GENE 1

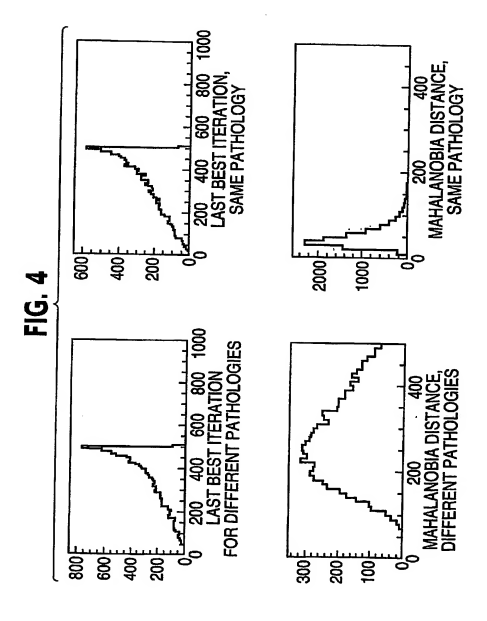
GENE 2

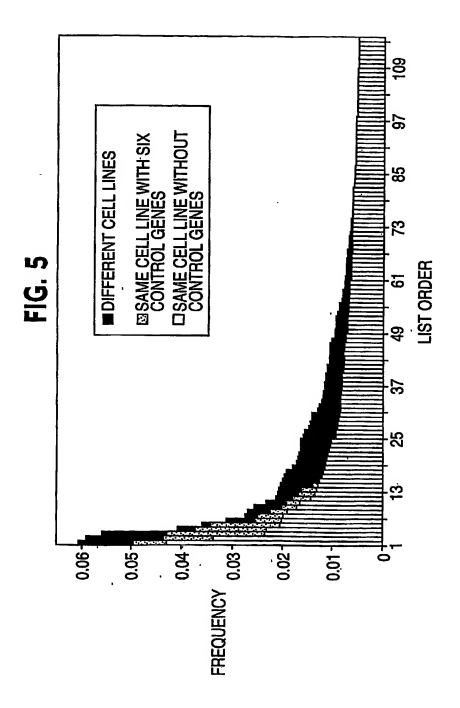
GENE K

GENE K

GENE C

GEN





# INTERNATIONAL SEARCH REPORT

International application No.

PC	T/	US	03/	05	73

T					
A. CLASSIFICATION OF SUBJECT MATTER  IPC(7) : C12Q 1/68; G06F 19/00  US CL : 435/6; 702/27					
According to International Patent Classification (IPC) or to both  B. FIELDS SEARCHED	national classification and IPC				
Minimum documentation searched (classification system follow	ed by classification symbols)				
U.S. : 435/6; 702/27					
Documentation searched other than minimum documentation to	the extent that such documents are included	d in the fields searched			
Electronic data base consulted during the international search (n	ame of data base and, where practicable, s	earch terms used)			
C. DOCUMENTS CONSIDERED TO BE RELEVANT					
Category *   Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to claim No.			
X Chen et al., Global Analysis of Gene Expression in Cancer Research, 01 July 2001. Volume 61, page	Chen et al., Global Analysis of Gene Expression in Invasion by a Lung Cancer Model, Cancer Research, 01 July 2001. Volume 61, pages 5223-5230, especially see Abstract; page 5224, column 1, Statistical Analysis section, column 2 to page 5225, Results				
A Wigle et al., Mining Mouse microarray data, Gen pages 1-5.	Wigle et al., Mining Mouse microarray data, Genome Biology, 2001, Volume 2, No. 7,				
Purther documents are listed in the continuation of Box C.					
* Special categories of olded documents:	See patent family annex.  The later document published after the inter-				
"A" document defining the general state of the art which is not considered to be of particular relevance	date and not in conflict with the application but cited to under principle or theory underlying the invention				
"E" carlier application or patent published on or after the international fling date	"X" document of particular relevance; the cl considered novel or cannot be considere when the document is taken alone	laimed invention cannot be ed to involve an inventive step			
*L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is				
"O" document referring to an oral disclosure, use, exhibition or other means	combined with one or more other such a being obvious to a person skilled in the	documents, such combination			
"P" document published prior to the international filing date but later than the priority date claimed	"&" document member of the same patent fa	mily			
Date of the actual completion of the international search	Date of mailing of the international search report				
04 November 2003 (04.11.2003)	0 9 JUN 2004				
Name and mailing address of the ISA/US  Mail Stop PCT, Atm: ISA/US  Commissioner for Patents	Authorized officer  Cheyne D Ly  Authorized Officer  Cheyne D Ly  Authorized Officer				
P.O. Box 1450 Alexandria, Virginia 22313-1450	Telephone No. 703 308-0196	For			
Facsimile No. (703)305-3230 orm PCT/ISA/210 (second sheet) (July 1998)					

## INTERNATIONAL SEARCH REPORT

International a	ppiication	NO.	

PCT/US03/05730

Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)					
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:					
Claim Nos.:  because they relate to subject matter not required to be searched by this Authority, namely:					
2. Claim Nos.:  because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:					
Claim Nos.:     because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).					
Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)	$\neg$				
This International Searching Authority found multiple inventions in this international application, as follows: Please See Continuation Sheet					
<ol> <li>As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.</li> <li>As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.</li> <li>As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:</li> </ol>					
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-13, species biological states and lung tissues  Remark on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.					
	1				

Form PCT/ISA/210 (continuation of first sheet(1)) (July 1998)



PCT/US03/05730

## BOX II. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING

This application contains claims directed to more than one species of the generic invention. These species are deemed to lack unity of invention because they are not so linked as to form a single general inventive concept under PCT Rule 13.1.

In order for more than one species to be examined, the appropriate additional examination fees must be paid. The species are as follows:

The instant invention is directed to 3 set of species listed in claims 2 (4 second states), claim 4 (8 tissue types), and claim 6 (8 cell types).

The species listed above do not relate to a single general inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, the species lack the same or corresponding special technical features for the following reasons:

According to PCT Rule 13.2 and to the guidelines in Section (f)(i)(A) of Annex B of the PCT Administrative Instructions, all alternatives of a Markush Group must have a common property or activity. Although the species of states, tissues, or cells of claims share a common feature of being either states, tissues, or cells; however, the species within each species set are not regarded as being of similar nature because all the alternatives do not share a common property or activity. For the species set of states in claim 2, the states do not share a common special technical feature because each state is defined by its mechanism of action and components involved. For the species set of tissues in claim 3 or cells in claim 6, the tissues or cell types do not share a common special technical feature because each state is defined by its mechanism of action and components involved. Therefore, the mechanisms and constituents of each species set of states, tissue types or cell types do not provide for a common property or activity or a significant structural element.

The instant invention is directed to 256 species. The first species are biological state, normal lung tissues, and normal lung cells. For each additional species per group thereafter, the fee would be \$210.00. Then, the cost for all for all of the species would be \$53,550.00.

Form PCT/ISA/210 (second sheet) (July 1998)

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